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(L12 AND 1BETA).USPT,PGPB,JPAB,EPAB,DWPI,TDBD.	10

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L14

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<i>DB=USPT,PGPB,JPAB,EPAB,DWPI,TDBD; PLUR=YES; OP=ADJ</i>			
<u>L14</u>	L12 and 1beta	10	<u>L14</u>
<u>L13</u>	L12 and beta1	1	<u>L13</u>
<u>L12</u>	L11 and beta	78	<u>L12</u>
<u>L11</u>	l3 and cancer	138	<u>L11</u>
<u>L10</u>	l3 cancer	7	<u>L10</u>
<u>L9</u>	wang-li-ming.in.	15	<u>L9</u>
<u>L8</u>	shelness-grgory.in.	0	<u>L8</u>
<u>L7</u>	childers-steven.in.	1	<u>L7</u>
<u>L6</u>	mach-robert-h.in.	3	<u>L6</u>
<u>L5</u>	wheeler-kenneth-t.in.	2	<u>L5</u>
<u>L4</u>	sigma-1beta	0	<u>L4</u>
<u>L3</u>	sigma\$5 receptor	650	<u>L3</u>
<u>L2</u>	sigma1beta	1	<u>L2</u>
<u>L1</u>	sigma1beta receptor	1	<u>L1</u>

END OF SEARCH HISTORY

STIC-Biotech/ChemLib

108586

From: Basi, Nirmal  
Sent: Tuesday, November 18, 2003 2:57 PM  
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Subject: sequence search for 09/823,069

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App. #: 09/832,069

Result format: Paper.

Title: Methods and composition utilizing an alternative splice variant of sigma-1 receptor

Inventors: Wheeler et al

Priority Date: 4/3/200

Please search:

i) SEQ ID NO: 1, 2

Search commercial and issued database.

Thanks,  
Nirmal S. Basi

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Online time: \_\_\_\_\_

TYPE OF SEARCH:  
NA Sequences: \_\_\_\_\_  
AA Sequences: \_\_\_\_\_  
Structures: \_\_\_\_\_  
Bibliographic: \_\_\_\_\_  
Litigation: \_\_\_\_\_  
Full text: \_\_\_\_\_  
Patent Family: \_\_\_\_\_  
Other: \_\_\_\_\_

VENDOR/COST (where applic.)  
STN: \_\_\_\_\_  
DIALOG: \_\_\_\_\_  
Questel/Orbit: \_\_\_\_\_  
DRLink: \_\_\_\_\_  
Lexis/Nexis: \_\_\_\_\_  
Sequence Sys.: \_\_\_\_\_  
WWW/Internet: \_\_\_\_\_  
Other (specify): \_\_\_\_\_

FILE 'MEDLINE'  
FILE 'JAPIO'  
FILE 'BIOSIS'  
FILE 'SCISEARCH'  
FILE 'WPIDS'  
FILE 'CAPLUS'  
FILE 'EMBASE'  
=> s signalbeta receptor#  
L1 1 SIGMA1BETA RECEPTOR#  
  
=> s signal receptor#  
L2 384 SIGMA1 RECEPTOR#  
  
=> s l2 and beta  
L3 21 L2 AND BETA  
  
=> dup rem l3  
PROCESSING COMPLETED FOR L3  
L4 14 DUP REM L3 (7 DUPLICATES REMOVED)  
  
=> d l1  
  
L1 ANSWER 1 OF 1 WPIDS COPYRIGHT 2003 THOMSON  
DERWENT ON STN  
AN 2001-662943 [76] WPIDS  
DNC C2001-194735  
TI Novel isolated polynucleotide encoding \*\*\*signalbeta\*\*\*  
\*\*\*receptor\*\*\* useful in screening assay to identify ligands specific  
for the \*\*\*signalbeta\*\*\* \*\*\*receptor\*\*\* for tumor imaging,  
diagnostic and treatment methods.  
DC B04 D16  
IN CHILDERS, S; MACH, R H; SHELNESS, G; WANG, L;  
WHEELER, K T  
PA (UYWA-N) UNIV WAKE FOREST; (CHIL-I) CHILDERS S;  
(MACH-I) MACH R H;  
(SHEL-I) SHELNESS G; (WANG-I) WANG L; (WHEE-I)  
WHEELER K T  
CYC 95  
PI WO 2001074297 A2 20011011 (200176)\* EN 56p A61K000-00  
RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT  
KE LS LU MC MW MZ  
NL OA PT SD SE SL SZ TR TZ UG ZW  
W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN  
CO CR CU CZ DE DK  
DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE  
KG KP KR KZ  
LK LC LR LS LT LU LV MA MD MG MK MN MW MX MZ  
NO NZ PL PT RO RU SD  
SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA  
ZW  
AU 2001087287 A 20011015 (200209) A61K000-00  
US 2002061847 A1 20020523 (200239) A61K038-17  
ADT WO 2001074297 A2 WO 2001-US10650 20010330; AU  
2001087287 A AU 2001-87287  
20010330; US 2002061847 A1 Provisional US 2000-193694P  
20000331, US  
2001-823069 20010330  
FDT AU 2001087287 A Based on WO 2001074297  
PRAI US 2000-193694P 20000331; US 2001-823069 20010330  
IC ICM A61K000-00; A61K038-17  
ICS C07H021-04; C07K014-705; C12N005-06; C12P021-02  
  
=> d ibib abs l3 1-21  
  
L3 ANSWER 1 OF 21 MEDLINE on STN  
ACCESSION NUMBER: 2000028702 MEDLINE  
DOCUMENT NUMBER: 20028702 PubMed ID: 10562962  
TITLE: Anti-amnesic effects of sigma (sigma)-receptor agonists.  
AUTHOR: Matsuno K  
CORPORATE SOURCE: Discovery Research Division, Santen  
Pharmaceutical Co.,  
Ltd., Osaka, Japan.  
SOURCE: NIPPON YAKURIGAKU ZASSHI. FOLIA  
PHARMACOLOGICA JAPONICA,  
(1999 Jul) 114 (1) 25-33. Ref. 50  
Journal code: 0420550. ISSN: 0015-5691.  
PUB. COUNTRY: Japan  
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
General Review; (REVIEW)  
(REVIEW, TUTORIAL)  
LANGUAGE: Japanese  
FILE SEGMENT: Priority Journals  
ENTRY MONTH: 200001  
ENTRY DATE: Entered STN: 20000204  
Last Updated on STN: 20000204  
Entered Medline: 20000121  
AB Both traditional and novel sigma (sigma)-receptor agonists have been  
reported to possess anti-amnesic effects in rodents. In particular, the  
anti-amnesic effects induced by the novel \*\*\*signal\*\*\* -  
\*\*\*receptor\*\*\* agonists, such as (+)-pentazocine, SA4503 and  
PRE-084,  
were shown in \*\*\*beta\*\*\* amyloid-peptide-induced, basal forebrain  
(BF)-lesioned and carbon monoxide (CO)-induced amnesia models and  
senescence-accelerated mouse (SAM). In addition, these  
\*\*\*signal\*\*\* -  
\*\*\*receptor\*\*\* agonists have good profiles for the central  
acetylcholine  
and dopamine systems. Moreover, they also have neuroprotective and  
anti-depressive effects. These evidence suggested that the  
\*\*\*signal\*\*\* -  
\*\*\*receptor\*\*\* agonists may be promising compounds for the  
treatment  
of dementing disorders such as Alzheimer's disease, senile dementia and  
vascular dementia. However, the sigma-receptor family is still  
considered

to be enigmatic molecular targets. More molecular cloning and  
biochemical  
studies on the sigma-receptor family are needed.  
  
L3 ANSWER 2 OF 21 MEDLINE on STN  
ACCESSION NUMBER: 1999330295 MEDLINE  
DOCUMENT NUMBER: 99330295 PubMed ID: 10403501  
TITLE: Ligands for opioid and sigma-receptors improve cardiac  
electrical stability in rat models of post-infarction  
cardiosclerosis and stress.  
AUTHOR: Lishmanov YuB; Maslov L N; Naryzhnaya N V; Tam S  
W  
CORPORATE SOURCE: Department of Experimental Cardiology,  
Institute of  
Cardiology, Tomsk, Russia.  
SOURCE: LIFE SCIENCES, (1999) 65 (1) PL13-7.  
Journal code: 0375521. ISSN: 0024-3205.  
PUB. COUNTRY: ENGLAND: United Kingdom  
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)  
LANGUAGE: English  
FILE SEGMENT: Priority Journals; Space Life Sciences  
ENTRY MONTH: 199907  
ENTRY DATE: Entered STN: 19990806  
Last Updated on STN: 19990806  
Entered Medline: 19990723  
AB The effects of the extremely selective mu-opioid receptor agonist,  
[D-Arg2,Lys4]-dormorphin-(1-4)-amide (DALDA), the mu-opioid  
receptor  
agonist morphine, the mu/delta agonist D-Ala2, Leu5, Arg6-enkephalin  
(dalgargin), the kappa-opioid receptor agonist spiradoline, and the  
\*\*\*signal\*\*\* - \*\*\*receptor\*\*\* antagonist DuP 734 on ventricular  
fibrillation threshold (VFT) was investigated in an experimental  
post-infarction cardiosclerosis model and an immobilization  
stress-induced  
model in rats. Both models produced a significant decrease in VFT.  
The  
postinfarction cardiosclerosis-induced decrease in VFT was significantly  
reversed by intravenous administration of dalgargin (0.1 mg/kg),  
DALDA (0.1  
mg/kg), or morphine HCl (1.5 mg/kg). Pretreatment with naloxone  
(0.2  
mg/kg) completely eliminated the increase in cardiac electrical stability  
produced by DALDA. Both spiradoline (8 mg/kg, i.p.) and DuP 734 (1  
mg/kg,  
i.p.) produced a significant increase in VFT in rats with post-infarction  
cardiosclerosis. This effect of spiradoline was blocked by  
nor-binaltorphimine. The immobilization stress-induced decrease in  
VFT  
was significantly reversed by administration of either DALDA,  
spiradoline  
or DuP 734. In conclusion, activation of either mu- or kappa1-opioid  
receptors or blockade of \*\*\*signal\*\*\* - \*\*\*receptors\*\*\* reversed the  
decrease in VFT in both cardiac compromised models. Since DALDA  
and  
dalgargin essentially do not cross blood brain barriers, their effects on  
VFT may be mediated through peripheral mu-opioid receptors.  
  
L3 ANSWER 3 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 2003:424096 BIOSIS  
DOCUMENT NUMBER: PREV200300424096  
TITLE: Cloning of an emopamil-binding protein (EBP)-like  
protein  
that lacks sterol DELTA8-DELTA7 isomerase activity.  
AUTHOR(S): Moebius, Fabian F. [Reprint Author]; Fitzky, Barbara  
U.;  
Wietzorek, Georg; Haidekker, Alexander; Eder, Andrea;  
Glossmann, Hartmut  
CORPORATE SOURCE: Institut fuer Biochemische Pharmakologie,  
Peter-Mayr-Strasse 1, A-6020, Innsbruck, Austria  
Fabian.Moebius@uibk.ac.at  
SOURCE: Biochemical Journal, (15 August, 2003) Vol. 374, No.  
1, pp.  
229-237. print.  
ISSN: 0264-6021.  
DOCUMENT TYPE: Article  
LANGUAGE: English  
OTHER SOURCE: DDBJ-AF034544; EMBL-AF034544;  
GenBank-AF034544;  
DDBJ-AF243433; EMBL-AF243433; GenBank-AF243433;  
DDBJ-U795328; EMBL-U795328; GenBank-U795328;  
DDBJ-Z37986;  
EMBL-Z37986; GenBank-Z37986  
ENTRY DATE: Entered STN: 17 Sep 2003  
Last Updated on STN: 17 Sep 2003  
AB EBP (emopamil-binding protein) is a high-affinity binding protein for  
(3H)emopamil and belongs to the family of so-called sigma receptors.  
Mutations that disrupt EBP's 3beta-hydroxysteroid sterol  
DELTA8-DELTA7  
isomerase activity (EC 5.3.3.5) impair cholesterol biosynthesis and  
cause  
X-chromosomal dominant chondrodysplasia punctata. We identified a  
human  
cDNA for a novel EBPL (EBP-like protein) with a calculated mass of  
23.2  
kDa. Amino acid sequence alignments and phylogenetic analysis  
revealed  
that EBPL is distantly related to EBP (31% identity and 52% similarity)  
and found in animals but not in plants. EBPL is encoded by four exons  
on  
human chromosome 13q14.2 covering 30.7 kb, and a partially  
processed EBPL  
pseudogene was found on 16q21. The EBPL mRNA was expressed  
ubiquitously  
and most abundant in liver, lung and kidney. Upon heterologous  
expression  
in yeast EBPL had no detectable 3beta-hydroxysteroid sterol

DELTA8-DELTA7  
isomerase and sigma-ligand-binding activity. Nine out of ten amino acid  
residues essential for catalytic activity of EBP were conserved in EBPL.  
Replacement of the only differing residue (EBP-Y111W) reduced  
catalytic  
activity of EBP. Transfer of the divergent residue from EBP to EBPL  
(EBPL-W91Y) and chimaerization of EBP and EBPL at various  
positions failed  
to restore catalytic activity of EBPL. Chemical cross-linking induced  
homodimerization of EBPL and EBP. Whereas mevinolin increased the  
mRNA  
for EBP and DHCR7 (DELTA7-sterol reductase) in HepG2 cells, it had  
no  
effect on mRNAs for EBPL and \*\*\*signal\*\*\* \*\*\*receptor\*\*\* ,  
indicating that EBP and EBPL expression are not coordinated. We  
propose  
that EBPL has a yet-to-be-discovered function other than cholesterol  
biosynthesis.  
  
L3 ANSWER 4 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 2003:223620 BIOSIS  
DOCUMENT NUMBER: PREV200300223620  
TITLE: Anti-amnesic effect of dimemorfan in mice.  
AUTHOR(S): Wang, Hui-Hung; Chien, Jyh-Wei; Chou,  
Yueh-Ching; Liao,  
Jyh-Fei [Reprint Author]; Chen, Chieh-Fu  
CORPORATE SOURCE: Department and Institute of Pharmacology,  
National  
Yang-Ming University, No. 155, Sec. 2, Li-Nong Street,  
Taipei, 112, Taiwan  
jfiao@ym.edu.tw  
SOURCE: British Journal of Pharmacology, (March 2003) Vol.  
138, No.  
5, pp. 941-949. print.  
ISSN: 0007-1188 (ISSN print).  
DOCUMENT TYPE: Article  
LANGUAGE: English  
ENTRY DATE: Entered STN: 7 May 2003  
Last Updated on STN: 7 May 2003  
AB 1. Dimemorfan, an antitussive for more than 25 years, has previously  
been  
reported to be a relative high-affinity ligand at sigma-1 ( \*\*\*signal\*\*\*  
) \*\*\*receptor\*\*\* with the Ki value of 151 nM. 2. To test whether  
dimemorfan has anti-amnesic effects similar to a \*\*\*signal\*\*\*  
\*\*\*receptor\*\*\* agonist, this study examined its effects on  
scopolamine-  
and \*\*\*beta\*\*\* -amyloid peptide-(25-35)-induced amnesia in mice. 3  
Dimemorfan (10-40 mg kg-1, i.p.) administered 30 min before the  
training  
trial, immediately after the training trial, or 30 min before the  
retention test significantly improved scopolamine (1 mg kg-1, i.p.)- or  
\*\*\*beta\*\*\* -amyloid peptide-(25-35) (3 nmol mouse-1,  
i.e.v.)-induced  
amnesia in a step-through passive avoidance test. Dimemorfan (5-40  
mg  
kg-1, i.p.) pretreatment also attenuated scopolamine (8 mg kg-1,  
i.p.)-induced amnesia in a water-maze test. And, these anti-amnesic  
effects of dimemorfan, like the putative \*\*\*signal\*\*\*  
\*\*\*receptor\*\*\*  
agonist (+)-N-allylnormetazocine ((+)-SKF-10047), were antagonized  
by a  
sigma receptor antagonist haloperidol (0.25 mg kg-1, i.p.). 4. These  
results indicated that dimemorfan has anti-amnesic effects and acts like  
a  
\*\*\*signal\*\*\* \*\*\*receptor\*\*\* agonist.  
  
L3 ANSWER 5 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 2002:551466 BIOSIS  
DOCUMENT NUMBER: PREV200200551466  
TITLE: Enhanced antidepressant effect of signal ( \*\*\*signal\*\*\* )  
\*\*\*receptor\*\*\* agonists in beta25-35-amyloid  
peptide-treated mice.  
AUTHOR(S): Urani, Alexandre; Romieu, Pascal; Roman, Francois  
J.;  
Maurice, Tangui [Reprint author]  
CORPORATE SOURCE: CNRS UMR 5102, University of Montpellier  
II, Place Eugene  
Bataillon, 34095, C.C. 090, Montpellier Cedex 5, France  
maurice@univ-montp2.fr  
SOURCE: Behavioural Brain Research, (21st August, 2002) Vol.  
134,  
No. 1-2, pp. 239-247. print.  
CODEN: BBREDI. ISSN: 0166-4328.  
DOCUMENT TYPE: Article  
LANGUAGE: English  
ENTRY DATE: Entered STN: 23 Oct 2002  
Last Updated on STN: 23 Oct 2002  
AB This study examined the antidepressant efficacy of the selective  
\*\*\*signal\*\*\* \*\*\*receptor\*\*\* agonists igmesine or PRE-084 in  
mice  
injected intracerebroventricularly (i.e.v.) with beta25-35-amyloid  
peptide  
and submitted to the forced swim test. beta25-35 peptide-injected  
animals  
developed memory deficits after 8 days contrarily to controls injected  
with scrambled beta25-35 peptide or vehicle solution. In the forced  
swim  
test, the i.e.v. treatment failed to affect the immobility duration, but  
the antidepressant effect of the signal agonists was facilitated in  
beta25-35 animals. Igmesine reduced immobility duration at 30 versus  
60  
mg/kg in control groups. PRE-084 decreased immobility duration at 30  
and  
60 mg/kg only in beta25-35 animals. Desipramine reduced the

immobility  
duration similarly among groups and fluoxetine appeared less potent in  
beta25-35 animals. The beta25-35 animals exhibited decreased  
progesterone  
levels in the hippocampus (~47%). The behavioural efficacy of sigma1  
agonists is known to depend on neuro(active)steroids levels synthesised  
by  
glial cells and neurones, which are affected by the \*\*\*beta\*\*\*  
-amyloid  
toxicity. This behavioural study suggests that sigma1 agonists, due to  
their enhanced efficacy, may allow to alleviate the depressive symptoms  
associated with Alzheimer's disease.

L3 ANSWER 6 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 2001:575419 BIOSIS  
DOCUMENT NUMBER: PREV200100575419  
TITLE: Enhanced antidepressant effect of sigma1 (sigma1)  
agonists  
in \*\*\*beta\*\*\* -amyloid peptide-treated rodents.  
AUTHOR(S): Urani, A. [Reprint author]; Romieu, P.; Roman, F. J.  
[Reprint author]; Noda, Y.; Kamei, H.; Tran, M. H.; Nagai,  
T.; Nabeshima, T.; Maurice, T.  
CORPORATE SOURCE: Biochimie/Enzymologie, Pfizer GRD, Fresnes,  
France  
SOURCE: Society for Neuroscience Abstracts, (2001) Vol. 27,  
No. 1,

pp. 853. print.  
Meeting Info.: 31st Annual Meeting of the Society for  
Neuroscience. San Diego, California, USA. November 10-15,  
2001.  
ISSN: 0190-5295.  
DOCUMENT TYPE: Conference; (Meeting)  
Conference; Abstract; (Meeting Abstract)  
LANGUAGE: English  
ENTRY DATE: Entered STN: 12 Dec 2001  
Last Updated on STN: 25 Feb 2002

AB The \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* is a 223 amino acid protein  
involved in numerous behavioral effects. In particular, \*\*\*sigma1\*\*\*  
\*\*\*receptor\*\*\* agonists present potent antidepressant-like effects in  
several animal models of behavioral despair. The antidepressant  
efficacy  
of selective sigma1 agonists was studied in two models of \*\*\*beta\*\*\*  
-amyloid-induced cognitive deficits. First, in mice injected centrally  
with beta25-35-amyloid peptide and submitted ten days after to forced  
swim  
test. In this test, igmesine appeared more efficient in beta25-35  
animals, by reducing immobility at 30 mg/kg vs. 60 mg/kg in control  
groups. Such facilitation was not observed with desipramine.  
Furthermore, beta25-35 animals exhibited decreased progesterone levels  
in  
the hippocampus (~47%). Second, in rats infused during 14 days with  
the  
beta1-40 amyloid peptide and submitted to the conditioned fear stress.  
In  
this test, (+)-SKF-10,047 reduced the stress-induced motor suppression  
at  
3 mg/kg in beta1-40 peptide infused rats, vs. 6 mg/kg in beta40-1  
treated  
rats. Igmesine presented an effect at 10 mg/kg in beta1-40 infused rats  
vs. 30 mg/kg in control rats. Neurosteroid measurements and  
immunohistochemical studies will also be presented. The sigma1  
agonist  
efficacy is known to depend on neuro(active)steroids levels, synthesized  
mainly by glial cells. These cells may be affected by b-amyloid toxicity.  
We suggest that sigma1 agonists, due to their enhanced efficacy, may  
improve Alzheimer's disease-related cognitive deficits.

L3 ANSWER 7 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 2001:503651 BIOSIS  
DOCUMENT NUMBER: PREV200100503651  
TITLE: (11C)Raclopride binding was reduced in vivo by  
\*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* ligand SA4503 in the  
mouse brain, while (11C)SA4503 binding was not by  
raclopride.  
AUTHOR(S): Ishiwa, Kiichi [Reprint author]; Kobayashi,  
Tadayuki;  
Kawamura, Kazunori; Matsuno, Kiyoshi; Senda, Michio  
CORPORATE SOURCE: Positron Medical Center, Tokyo Metropolitan  
Institute of  
Gerontology, Tokyo, Japan  
ishiwa@pet.imig.or.jp  
SOURCE: Nuclear Medicine and Biology, (October, 2001) Vol.  
28, No.  
7, pp. 787-792. print.  
ISSN: 0969-8051.

DOCUMENT TYPE: Article  
LANGUAGE: English  
ENTRY DATE: Entered STN: 31 Oct 2001  
Last Updated on STN: 23 Feb 2002

AB (11C)Raclopride is widely used as a representative dopamine D2-like  
receptor ligand in positron emission tomography (PET) studies, and  
(11C)1-(3,4-dimethoxyphenethyl)-4-(3-phenylpropyl)piperazine  
dihydrochloride (11C)SA4503 is a recently developed selective ligand  
for  
mapping \*\*\*sigma1\*\*\* \*\*\*receptors\*\*\* in the brain. The striatal  
uptake of (11C)raclopride in mice was reduced by co-injection of an  
excess  
amount of SA4503, in spite of the fact that raclopride had no effect on  
the brain uptake of (11C)SA4503 as shown in a previous study. The  
blocking effect of SA4503 on the striatal uptake of (11C)raclopride was  
dose-dependent, but disappeared by 1 h or 6 h after intraperitoneal  
injection of SA4503. The brain uptake of (11C)SA4503 was not  
affected by  
a dopamine transporter inhibitor GBR 12909, nor was (11C)  
\*\*\*beta\*\*\*  
-CIT-FP inhibited by SA4503. The IC50 values of raclopride for

signal and  
sigma2 receptor subtypes measured in vitro were 11800 nM and 4950  
nM,  
respectively, suggesting that the affinity was too low for  
(11C)raclopride  
to bind in vivo to sigma receptors. On the other hand, the IC50 value  
of  
SA4503 for dopamine D2 receptors was 470 nM, that is approximate  
1/25 of  
the affinity of raclopride for the dopamine D2 receptors. Therefore,  
possible explanations for the partial blocking effects of SA4503 on the  
striatal uptake of (11C)raclopride are: (1) an excess amount of SA4503  
may  
reduce the (11C)raclopride uptake due to its low affinity for dopamine  
D2  
receptors, or (2) SA4503 may enhance endogenous dopamine release,  
which  
results in the competitive inhibition of the (11C)raclopride uptake.  
These findings support that both (11C)raclopride and (11C)SA4503 are  
selective in vivo ligands for dopamine D2-like receptors and  
\*\*\*sigma1\*\*\* \*\*\*receptors\*\*\*, respectively, in spite of the  
partial  
blocking effect of SA4503 on the striatal uptake of (11C)raclopride.

L3 ANSWER 8 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 1999:477414 BIOSIS  
DOCUMENT NUMBER: PREV199900477414  
TITLE: Anti-amnesic effects of sigma (sigma) -receptor agonists.  
AUTHOR(S): Matsuno, Kiyoshi [Reprint author]  
CORPORATE SOURCE: Discovery Research Division, Santen  
Pharmaceutical Co.,  
Ltd., Shimoshinoji, Higashiyodogawa, Osaka, 533-8651, Japan  
SOURCE: Folia Pharmacologica Japonica, (July, 1999) Vol. 114,  
No.  
1, pp. 25-33. print.  
CODEN: NYKZAU. ISSN: 0015-5691.

DOCUMENT TYPE: Article  
LANGUAGE: Japanese  
ENTRY DATE: Entered STN: 9 Nov 1999  
Last Updated on STN: 9 Nov 1999  
AB Both traditional and novel sigma (sigma) -receptor agonists have been  
reported to possess anti-amnesic effects in rodents. In particular, the  
anti-amnesic effects induced by the novel \*\*\*sigma1\*\*\* -  
\*\*\*receptor\*\*\* agonists, such as (+) -pentazocine, SA4503 and  
PRE-084,  
were shown in \*\*\*beta\*\*\* amyloid-peptide-induced, basal forebrain  
(BF)  
- lesioned and carbon monoxide (CO) -induced amnesia models and  
senescence-accelerated mouse (SAM). In a dition, these  
\*\*\*sigma1\*\*\* -  
\*\*\*receptor\*\*\* agonists have good profiles for the central  
acetylcholine  
and dopamine systems. Moreover, they also have neuroprotective and  
anti-depressive effects. These evidence suggested that the  
\*\*\*sigma1\*\*\*  
- \*\*\*receptor\*\*\* agonists may be promising compounds for the  
treatment  
of dementing disorders such as Alzheimer's disease, senile dementia and  
vascular dementia. However, the sigma-receptor family is still  
considered  
to be enigmatic molecular targets. More molecular cloning and  
biochemical  
studies on the sigma-receptor family are needed.

L3 ANSWER 9 OF 21 BIOSIS COPYRIGHT 2003 BIOLOGICAL  
ABSTRACTS INC. on STN  
ACCESSION NUMBER: 1998:98297 BIOSIS  
DOCUMENT NUMBER: PREV199800098297  
TITLE: Sigma1 ( \*\*\*sigma1\*\*\* ) \*\*\*receptor\*\*\* agonists  
and  
neurosteroids attenuate B22-35-amyloid peptide-induced  
amnesia in mice through a common mechanism.  
AUTHOR(S): Maurice, T. [Reprint author]; Su, T.-P.; Privat, A.  
CORPORATE SOURCE: I.N.S.E.R.M. Unite 336, Dev. Plasticite,  
Vieillessement du  
Systeme Nerveux, 8 rue de l'Ecole Normale, 34296  
Montpellier Cedex 5, France  
SOURCE: Neuroscience, (March, 1998) Vol. 83, No. 2, pp.  
413-428.  
print.  
CODEN: NRSCDN. ISSN: 0306-4522.

DOCUMENT TYPE: Article  
LANGUAGE: English  
ENTRY DATE: Entered STN: 25 Feb 1998  
Last Updated on STN: 6 Apr 1998  
AB The sigma1 ( \*\*\*sigma1\*\*\* ) \*\*\*receptor\*\*\* agonists exert  
potent  
anti-amnesic effects, as they apparently block the learning impairments  
either induced by the muscarinic receptor antagonist scopolamine, the  
N-methyl-D-aspartate receptor antagonist dizocilpine or inherently due  
to  
the age-related deficits in senescence-accelerated mice. We recently  
described the amnesia induced by the \*\*\*beta\*\*\* -amyloid-related  
peptide beta25-35, administered centrally in an aggregated form, in  
mice.  
The deficits were sensitive to cholinomimetics or to N-methyl-D-  
aspartate/glycine modulatory site agonists. Herein, we examined the  
effects of \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* ligands on the beta25-35  
peptide-induced amnesia. The effects of neuro(active) steroids, which  
interact in vitro and in vivo with \*\*\*sigma1\*\*\* \*\*\*receptors\*\*\*  
were examined in parallel. Mnesic capacity was evaluated seven days  
after  
administration of aggregated beta25-35 peptide (3 nmol), using  
spontaneous  
alternation in the Y-maze for spatial short-term memory, or after 14  
days,  
using the step-down type passive avoidance test for long-term memory.

The  
\*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* agonists (+)-pentazocine,  
PRE-084, or  
SA4503 attenuated, in a dose-dependent and bell-shaped manner, the  
beta25-35 peptide-induced deficits on both tests. These effects were  
antagonized by haloperidol or BMY-14802, confirming the  
\*\*\*sigma1\*\*\*  
\*\*\*receptor\*\*\* pharmacology. Pregnenolone,  
dehydroepiandrosterone, and  
their sulphate esters, but not progesterone, also dose-dependently  
attenuated the beta25-35 peptide-induced deficits. Progesterone  
blocked  
the beneficial effects of each other neurosteroid, behaving as an  
antagonist. Furthermore, haloperidol blocked the effects induced by  
neurosteroids, whereas progesterone antagonized the effects of the  
non-steroidal \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* agonists, showing a  
clear crossed pharmacology of different drug classes. These results  
demonstrated that: (i) the anti-amnesic effect of \*\*\*sigma1\*\*\*  
\*\*\*receptor\*\*\* agonists may be of therapeutic relevance in  
pathological  
states affecting the cholinergic and/or glutamatergic systems, such as in  
pathological aging; (ii) neurosteroids play an important role in learning  
processes and may collectively constitute a therapeutic target; (iii) the  
interaction between sigma1 systems and neurosteroids appears indeed  
of  
behavioural relevance.

L3 ANSWER 10 OF 21 CAPLUS COPYRIGHT 2003 ACS on STN  
ACCESSION NUMBER: 2003:619310 CAPLUS  
DOCUMENT NUMBER: 139:287042  
TITLE: Cloning of an emopamil-binding protein (EBP)-like  
protein that lacks sterol .DELTA.8-.DELTA.7 isomerase  
activity  
AUTHOR(S): Moebius, Fabian F.; Fitzky, Barbara U.;  
Wietzorrek,  
Georg; Haidekker, Alexander; Eder, Andrea; Glossmann,  
Hartmut  
CORPORATE SOURCE: Institut fuer Biochemische Pharmakologie,  
Innsbruck,  
A-6020, Austria  
SOURCE: Biochemical Journal (2003), 374(1), 229-237  
CODEN: BJOAK; ISSN: 0264-6021  
PUBLISHER: Portland Press Ltd.  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB EBP (emopamil-binding protein) is a high-affinity binding protein for  
[3H]emopamil and belongs to the family of so-called sigma receptors.  
Mutations that disrupt EBP's 3. \*\*\*beta\*\*\* -hydroxysteroid sterol  
.DELTA.8-.DELTA.7 isomerase activity (EC 5.3.3.5) impair  
cholesterol  
biosynthesis and cause X-chromosomal dominant chondrodysplasia  
punctata.

The authors identified a human cDNA for a novel EBPL (EBP-like  
protein)  
with a calcd. mass of 23.2 kDa. Amino acid sequence alignments and  
phylogenetic anal. revealed that EBPL is distantly related to EBP (31%  
identity and 52% similarity) and found in animals but not in plants.  
EBPL  
is encoded by four exons on human chromosome 13q14.2 covering 30.7  
kb, and  
a partially processed EBPL pseudogene was found on 16q21. The  
EBPL mRNA  
was expressed ubiquitously and most abundant in liver, lung and kidney  
Upon heterologous expression in yeast EBPL had no detectable 3.  
\*\*\*beta\*\*\* -hydroxysteroid sterol .DELTA.8-.DELTA.7 isomerase  
and  
sigma-ligand-binding activity. Nine out of ten amino acid residues  
essential for catalytic activity of EBP were conserved in EBPL.  
Replacement of the only differing residue (EBP-Y11W) reduced  
catalytic  
activity of EBP. Transfer of the divergent residue from EBP to EBPL  
(EBPL-W91Y) and chimerization of EBP and EBPL at various  
positions failed  
to restore catalytic activity of EBPL. Chem. crosslinking induced  
homodimerization of EBPL and EBP. Whereas mevinolin increased the  
mRNA  
for EBP and DHCR7 (.DELTA.7-sterol reductase) in HepG2 cells, it  
had no  
effect on mRNAs for EBPL and \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\*,  
indicating that EBP and EBPL expression are not coordinated. The  
authors  
propose that EBPL has a yet-to-be-discovered function other than  
cholesterol biosynthesis.  
REFERENCE COUNT: 30 THERE ARE 30 CITED  
REFERENCES AVAILABLE FOR THIS  
RECORD. ALL CITATIONS AVAILABLE IN THE

RE FORMAT

L3 ANSWER 11 OF 21 CAPLUS COPYRIGHT 2003 ACS on STN  
ACCESSION NUMBER: 2003:375472 CAPLUS  
DOCUMENT NUMBER: 139:160091  
TITLE: .sigma.1 Receptor-related neuroactive steroids  
modulate cocaine-induced reward  
AUTHOR(S): Romieu, Pascal; Martin-Fardon, Remi; Bowen,  
Wayne D.;  
Maurice, Tangui  
CORPORATE SOURCE: Centre National de la Recherche  
Scientifique Unite  
Mixte de Recherche 5102, University of Montpellier II,  
Montpellier, 34095/5, Fr.  
SOURCE: Journal of Neuroscience (2003), 23(9), 3572-3576  
CODEN: JNRSDS; ISSN: 0270-6474  
PUBLISHER: Society for Neuroscience  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB The .sigma.1 receptor is critically involved in the rewarding effect of  
cocaine, as measured using the conditioned place preference (CPP)  
procedure in mice. Neuroactive steroids exert rapid neuromodulatory

effects in the brain by interacting with GABAA, NMDA, and .sigma.1 receptors. At the .sigma.1 receptor level, 3. \*\*\*beta\*\*\* -hydroxy-5-androsten-17-one [dihydroepiandrosterone (DHEA)] and 3. \*\*\*beta\*\*\* -hydroxy-5-pregnen-20-one (pregnenolone) act as agonists, whereas 4-pregnen-3,20-dione (progesterone) is an efficient antagonist. The present study sought to investigate the action of neuroactive steroids in acquisition of cocaine-induced CPP in C57BL/6 mice. None of these steroids induced CPP alone. However, pretreatment with DHEA or pregnenolone (5-20 mg/kg, s.c.) during conditioning with cocaine (10 mg/kg, i.p.) increased the conditioned score. On the contrary, pretreatment with either progesterone (10 or 20 mg/kg, s.c.) or finasteride (25 mg/kg, twice a day), a 5.alpha.-reductase inhibitor, blocked acquisition of cocaine (20 mg/kg)-induced CPP. A crossed pharmacol. was obsd. between steroids and .sigma.1 ligands. The .sigma.1 antagonist N-[2-(3,4-dichlorophenyl)ethyl]-N-methyl-2-(dimethylamino)ethylamine blocked cocaine-induced CPP and its potentiation by DHEA or pregnenolone. Progesterone blocked cocaine-induced CPP and its potentiation by the .sigma.1 agonist igmesine. These results showed that neuroactive steroids play a role in cocaine-induced appetence, through their interaction with the .sigma.1 receptor. Therefore, neuroendocrine control of cocaine addiction may not involve solely glucocorticoids. The importance of neuroactive steroids as factors of individual vulnerability to drug addiction should, thus, be considered. REFERENCE COUNT: 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 12 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 2003:216650 CAPLUS  
DOCUMENT NUMBER: 138:336192  
TITLE: IL-10 Mediates \*\*\*Sigma1\*\*\* \*\*\*Receptor\*\*\*  
-Dependent Suppression of Antitumor Immunity  
AUTHOR(S): Zhu, Li X.; Sharma, Sherven; Gardner, Brian; Escudro, Brian; Atianzar, Kimberly; Tashkin, Donald P.; Dubinett, Steven M.  
CORPORATE SOURCE: Division of Pulmonary and Critical Care Medicine, David Geffen School of Medicine, University of California, Los Angeles, CA, 90095, USA  
SOURCE: Journal of Immunology (2003), 170(7), 3585-3591  
CODEN: JOIMA3; ISSN: 0022-1767  
PUBLISHER: American Association of Immunologists  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB Sigma receptors are unique endoplasmic reticulum proteins that mediate signaling for a variety of drugs. The authors detd. the effect of \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* agonists on immune responses in a syngeneic lung cancer model. \*\*\*Sigma1\*\*\* \*\*\*receptor\*\*\* agonists, including cocaine, up-regulated splenocyte IL-10 mRNA and protein prodn. in vitro in a sigma receptor-dependent, pertussis toxin-sensitive manner. In vivo, \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* agonists promoted tumor growth and induced IL-10 at the tumor site. Increased tumor growth was prevented by administration of specific Abs to IL-10 or by administration of specific \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* antagonists. The authors report that \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* ligands, including cocaine, augment tumor growth through an IL-10 dependent mechanism. REFERENCE COUNT: 60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 13 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 2003:12042 CAPLUS  
DOCUMENT NUMBER: 138:248840  
TITLE: .sigma.1 receptor agonist-mediated regulation of N-methyl-D-aspartate-stimulated [3H]dopamine release is dependent upon protein kinase C  
AUTHOR(S): Nuwayhid, Samer J.; Werling, Linda L.  
CORPORATE SOURCE: Department of Pharmacology, The George Washington University Medical Center, Washington, DC, USA  
SOURCE: Journal of Pharmacology and Experimental Therapeutics (2003), 304(1), 364-369  
CODEN: JPETAB; ISSN: 0022-3265  
PUBLISHER: American Society for Pharmacology and Experimental Therapeutics  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB The authors have previously shown that .sigma.1 receptor agonists inhibit N-methyl-D-aspartate (NMDA)-stimulated [3H]dopamine from slices of striatum in a concn.-related manner and that the inhibition is reversed by .sigma.1 receptor-selective and nonsubtype-selective .sigma.1 receptor antagonists. Based on previous evidence from the authors' lab. as well as other labs., the authors hypothesized that .sigma.1 receptors might use a protein kinase C (PKC) signaling pathway to modulate stimulated dopamine release. The authors tested several inhibitors of PKC isoenzymes, as

well as a phospholipase C inhibitor for their effects on .sigma.1 receptor agonist-mediated regulation of [3H]dopamine release. Although none of the inhibitors tested affected the ability of NMDA to stimulate [3H]dopamine release, they all abolished regulation by the .sigma.1 receptor agonist (+)-pentazocine in a concn.-related manner. The authors also found that prior exposure to 1 .mu.M phorbol 2-myristate 13-acetate for 30 min abolished regulation by (+)-pentazocine. The authors concluded that an intact PKC system was required for .sigma.1 agonist-mediated regulation of NMDA-stimulated [3H]dopamine release from rat striatal slices. Based on the pharmacol. profile of the PKC inhibitors tested, as well as reports in the literature on PKC involvement in neurotransmitter release and .sigma.1 receptor action, PKC. \*\*\*beta\*\*\* seems most likely to be responsible, at least in part, for the effects of (+)-pentazocine on dopamine release. REFERENCE COUNT: 40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 14 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 2002:634631 CAPLUS  
DOCUMENT NUMBER: 138:348561  
TITLE: Enhanced antidepressant effect of sigma1 (.sigma.1) receptor agonists in . \*\*\*beta\*\*\* .25-35-amyloid peptide-treated mice  
AUTHOR(S): Urani, Alexandre; Romieu, Pascal; Roman, Francois J.; Maurice, Tanguj  
CORPORATE SOURCE: Behavioural Neuropharmacology Group, INSERM U.336, Institut de Biologie, Montpellier, 34060, Fr.  
SOURCE: Behavioural Brain Research (2002), 134(1,2), 239-247  
CODEN: BBREDI; ISSN: 0166-4328  
PUBLISHER: Elsevier Science B.V.  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB This study examd. the antidepressant efficacy of the selective .sigma.1 receptor agonists igmesine or PRE-084 in mice injected intracerebroventricularly (i.c.v.) with . \*\*\*beta\*\*\* .25-35-amyloid peptide and submitted to the forced swim test. . \*\*\*beta\*\*\* .25-35 Peptide-injected animals developed memory deficits after 8 days contrarily to controls injected with scrambled . \*\*\*beta\*\*\* .25-35 peptide or vehicle soln. In the forced swim test, the i.c.v. treatment failed to affect the immobility duration, but the antidepressant effect of the .sigma.1 agonists was facilitated in . \*\*\*beta\*\*\* .25-35 animals. Igmesine reduced immobility duration at 30 vs. 60 mg/kg in control groups. PRE-084 decreased immobility duration at 30 and 60 mg/kg only in . \*\*\*beta\*\*\* .25-35 animals. Desipramine reduced the immobility duration similarly among groups and fluoxetine appeared less potent in . \*\*\*beta\*\*\* .25-35 animals. The . \*\*\*beta\*\*\* .25-35 animals exhibited decreased progesterone levels in the hippocampus (-47%). The behavioral efficacy of .sigma.1 agonists is known to depend on neuro(active)steroids levels synthesized by glial cells and neurons, which are affected by the . \*\*\*beta\*\*\* -amyloid toxicity. This behavioral study suggests that .sigma.1 agonists, due to their enhanced efficacy, may allow to alleviate the depressive symptoms assocd. with Alzheimer's disease. REFERENCE COUNT: 44 THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 15 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 2001:709266 CAPLUS  
DOCUMENT NUMBER: 136:382245  
TITLE: [11C]Raclopride binding was reduced in vivo by \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* ligand SA4503 in the mouse brain, while [11C]SA4503 binding was not by raclopride  
AUTHOR(S): Ishiwa, K.; Kobayashi, T.; Kawamura, K.; Matsuno, K.; Senda, M.  
CORPORATE SOURCE: Positron Medical Center, Tokyo Metropolitan Institute of Gerontology, Tokyo, Japan  
SOURCE: Nuclear Medicine and Biology (2001), 28(7), 787-792  
CODEN: NMBIEO; ISSN: 0969-8051  
PUBLISHER: Elsevier Science Inc.  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB [11C]Raclopride is widely used as a representative dopamine D2-like receptor ligand in positron emission tomog. (PET) studies, and [11C]1-(3,4-dimethoxyphenethyl)-4-(3-phenylpropyl)piperazine dihydrochloride ([11C]SA4503) is a recently developed selective ligand for mapping \*\*\*sigma1\*\*\* \*\*\*receptors\*\*\* in the brain. The striatal uptake of [11C]raclopride in mice was reduced by co-injection of an excess amt. of SA4503, in spite of the fact that raclopride had no effect on the brain uptake of [11C]SA4503 as shown in a previous study. The blocking effect of SA4503 on the striatal uptake of [11C]raclopride was dose-dependent, but disappeared by 1 h or 6 h after i.p. injection of SA4503. The brain uptake of [11C]SA4503 was not affected by a

dopamine transporter inhibitor GBR 12909, nor was [11C]. \*\*\*beta\*\*\* -CIT-FP inhibited by SA4503. The IC50 values of raclopride for sigma1 and sigma2 receptor subtypes measured in vitro were 11800 nM and 4950 nM, resp., suggesting that the affinity was too low for [11C]raclopride to bind in vivo to sigma receptors. On the other hand, the IC50 value of SA4503 for dopamine D2 receptors was 470 nM, that is approx. 1/25 of the affinity of raclopride for the dopamine D2 receptors. Therefore, possible explanations for the partial blocking effects of SA4503 on the striatal uptake of [11C]raclopride are: (1) an excess amt. of SA4503 may reduce the [11C]raclopride uptake due to its low affinity for dopamine D2 receptors, or (2) SA4503 may enhance endogenous dopamine release, which results in the competitive inhibition of the [11C]raclopride uptake. These finding support that both [11C]raclopride and [11C]SA4503 are selective in vivo ligands for dopamine D2-like receptors and \*\*\*sigma1\*\*\* \*\*\*receptors\*\*\*, resp., in spite of the partial blocking effect of SA4503 on the striatal uptake of [11C]raclopride. REFERENCE COUNT: 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 16 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 2000:255286 CAPLUS  
DOCUMENT NUMBER: 133:53926  
TITLE: Immunocytochemical localization of the \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* in the adult rat central nervous system  
AUTHOR(S): Alonso, G.; Phan, V.-L.; Guillemain, I.; Saunier, M.; Legrand, A.; Anoaï, M.; Maurice, T.  
CORPORATE SOURCE: INSERM Unite 336, Developpement, Plasticite et Vieillessement du Systeme Nerveux, Montpellier, Fr.  
SOURCE: Neuroscience (Oxford) (2000), 97(1), 155-170  
CODEN: NRSCDN; ISSN: 0306-4522  
PUBLISHER: Elsevier Science Ltd.  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB To characterize the localization of the \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* in the adult rat central nervous system, a polyclonal antibody was raised against a 20 amino acid peptide, corresponding to the fragment 143-162 of the cloned \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* protein. Throughout the rostrocaudal regions of the central nervous system extending from the olfactory bulb to the spinal cord, intense to moderate immunostaining was found to be assocd. with: (i) ependymocytes bordering the entire ventricular system, and (ii) neuron-like structures located within the parenchyma. Double fluorescence studies confirmed that, throughout the parenchyma, \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* -immunostaining was essentially assocd. with neuronal structures immunostained for the neuronal marker . \*\*\*beta\*\*\* .III-tubulin. In all rats examd., high levels of immunostaining were always assocd. with neurons located within specific regions including the granular layer of the olfactory bulb, various hypothalamic nuclei, the septum, the central gray, motor nuclei of the hindbrain and the dorsal horn of the spinal cord. In contrast, only faint immunostaining was assocd. with neurons located in the caudate-putamen and the cerebellum. Electron microscope studies indicated that \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* immunostaining was mostly assocd. with neuronal perikarya and dendrites, where it was localized to the limiting plasma membrane, the membrane of mitochondria and of some cisternae of the endoplasmic reticulum. At the level of synaptic contacts, intense immunostaining was assocd. with postsynaptic structures including the postsynaptic thickening and some polymorphous vesicles, whereas the presynaptic axons were devoid of immunostaining. These data indicate that the \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* antibody prepd. here, represents a promising tool for further investigating the role of .sigma.1 receptors. REFERENCE COUNT: 41 THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 17 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 1999:507828 CAPLUS  
DOCUMENT NUMBER: 131:255271  
TITLE: Intracellular .sigma.1 receptor modulates phospholipase C and protein kinase C activities in the brainstem  
AUTHOR(S): Morin-Surun, M. P.; Collin, T.; Denavit-Saubie, M.; Baulieu, E.-E.; Monnet, F. P.  
CORPORATE SOURCE: Institut Alfred Fessard, Gif-sur Yvette, 91198, Fr.  
SOURCE: Proceedings of the National Academy of Sciences of the United States of America (1999), 96(14), 8196-8199  
CODEN: PNASA6; ISSN: 0027-8424

PUBLISHER: National Academy of Sciences  
DOCUMENT TYPE: Journal  
LANGUAGE: English  
AB Most physiol. effects of .sigma.1 receptor ligands are sensitive to pertussis toxin, suggesting a coupling with cell membrane-bound G proteins. However, the cloning of the .sigma.1 receptor has allowed the identification of an intracellular protein anchored on the endoplasmic reticulum. Here, we show, using the isolated adult guinea pig brainstem prepn., that activation of the .sigma.1 receptor results in its translocation from the cytosol to the vicinity of the cell membrane and induces a robust and rapid decrease in hypoglossal activity, which is mediated by phospholipase C. The subsequent activation of protein kinase C. \*\*\*beta\*\*\* .1 and . \*\*\*beta\*\*\* .2 isoforms and the phosphorylation of a protein of the same mol. wt. as the cloned .sigma.1 receptor lead to a desensitization of the .sigma.1 motor response. Our results indicate that the intracellular .sigma.1 receptor regulates several components implicated in plasma membrane-bound signal transduction. This might be an example of a mechanism by which an intracellular receptor modulates metabotropic responses.  
REFERENCE COUNT: 25 THERE ARE 25 CITED  
REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 18 OF 21 CAPLUS COPYRIGHT 2003 ACS ON STN  
ACCESSION NUMBER: 1999:419391 CAPLUS  
DOCUMENT NUMBER: 131:96849  
TITLE: Anti-amnesic effects of sigma (.sigma.)-receptor agonists  
AUTHOR(S): Matsuno, Kiyoshi  
CORPORATE SOURCE: Discovery Res. Div., Santen Pharm. Co., Ltd., Osaka, 533-8651, Japan  
SOURCE: Nippon Yakurigaku Zasshi (1999), 114(1), 25-33  
CODEN: NYKZAU; ISSN: 0015-5691  
PUBLISHER: Nippon Yakuri Gakkai  
DOCUMENT TYPE: Journal; General Review  
LANGUAGE: Japanese  
AB A review with 50 refs. Both traditional and novel sigma (.sigma.)-receptor agonists have been reported to possess anti-amnesic effects in rodents. In particular, the anti-amnesic effects induced by the novel \*\*\*sigma1\*\*\* - \*\*\*receptor\*\*\* agonists, such as (+)-pentazocine, SA4503, and PRE-084, were shown in . \*\*\*beta\*\*\* amyloid-peptide-induced, basal forebrain (BF) lesioned and carbon monoxide (CO)-induced amnesia models and senescence-accelerated mouse (SAM). In addn., these \*\*\*sigma1\*\*\* - \*\*\*receptor\*\*\* agonists have good profiles for the central acetylcholine and dopamine systems. Moreover, they also have neuroprotective and anti-depressive effects. These evidence suggested that the \*\*\*sigma1\*\*\* - \*\*\*receptor\*\*\* agonists may be promising compds. for the treatment of dementing disorders such as Alzheimer's disease, senile dementia, and vascular dementia. However, the sigma-receptor family is still considered to be enigmatic mol. targets. More mol. cloning and biochem. studies on the sigma-receptor family are needed.

L3 ANSWER 19 OF 21 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN  
ACCESSION NUMBER: 2000131045 EMBASE  
TITLE: Immunocytochemical localization of the \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* in the adult rat central nervous system.  
AUTHOR: Alonso G.; Phan V.-L.; Guillemain I.; Saunier M.; Legrand A.; Anoaï M.; Maurice T.  
CORPORATE SOURCE: T. Maurice, Developpement, Vieillessement Systeme Nerveux, INSERM Unite 336, Montpellier, France  
SOURCE: Neuroscience, (2000) 97/1 (155-170).  
Refs: 41  
ISSN: 0306-4522 CODEN: NRSCDN  
PUBLISHER IDENT.: S 0306-4522(00)00014-2  
COUNTRY: United Kingdom  
DOCUMENT TYPE: Journal; Article  
FILE SEGMENT: 001 Anatomy, Anthropology, Embryology and Histology  
026 Immunology, Serology and Transplantation  
LANGUAGE: English  
SUMMARY LANGUAGE: English  
AB In order to characterize the localization of the \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* in the adult rat central nervous system, a polyclonal antibody was raised against a 20 amino acid peptide, corresponding to the fragment 143-162 of the cloned \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* protein. Throughout the rostrocaudal regions of the central nervous system extending from the olfactory bulb to the spinal cord, intense to moderate immunostaining was found to be associated with: (i) ependymocytes bordering the entire ventricular system, and (ii) neuron-like structures located within the parenchyma. Double fluorescence studies confirmed that, throughout the parenchyma, \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* -immunostaining was essentially associated with neuronal structures immunostained for the neuronal marker . \*\*\*beta\*\*\* .III-tubulin. In all rats examined, high levels of immunostaining were always associated with neurons located within specific regions including the granular layer of

the olfactory bulb, various hypothalamic nuclei, the septum, the central gray, motor nuclei of the hindbrain and the dorsal horn of the spinal cord. In contrast, only faint immunostaining was associated with neurons located in the caudate-putamen and the cerebellum. Electron microscope studies indicated that \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* immunostaining was mostly associated with neuronal perikarya and dendrites, where it was localized to the limiting plasma membrane, the membrane of mitochondria and of some cisternae of the endoplasmic reticulum. At the level of synaptic contacts, intense immunostaining was associated with postsynaptic structures including the postsynaptic thickening and some polymorphous vesicles, whereas the presynaptic axons were devoid of immunostaining. These data indicate that the \*\*\*sigma1\*\*\* \*\*\*receptor\*\*\* antibody prepared here, represents a promising tool for further investigating the role of .sigma.1 receptors. Copyright (C) 2000 IBRO.

L3 ANSWER 20 OF 21 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN  
ACCESSION NUMBER: 1999247997 EMBASE  
TITLE: Anti-amnesic effects of sigma (.sigma.) -receptor agonists.  
AUTHOR: Matsuno K.  
CORPORATE SOURCE: K. Matsuno, Discovery Research Division, Santen Pharmaceutical Co., Ltd., Shimoshinjo, Higashiyodogawa, Osaka 533-8651, Japan  
SOURCE: Folia Pharmacologica Japonica, (1999) 114/1 (25-33).  
Refs: 50  
ISSN: 0015-5691 CODEN: NYKZAU  
COUNTRY: Japan  
DOCUMENT TYPE: Journal; Article  
FILE SEGMENT: 008 Neurology and Neurosurgery  
030 Pharmacology  
037 Drug Literature Index  
LANGUAGE: Japanese  
SUMMARY LANGUAGE: English; Japanese  
AB Both traditional and novel sigma (.sigma.)-receptor agonists have been reported to possess anti-amnesic effects in rodents. In particular, the anti- amnesic effects induced by the novel \*\*\*sigma1\*\*\* - \*\*\*receptor\*\*\* agonists, such as (+)- pentazocine, SA4503 and PRE-084, were shown in . \*\*\*beta\*\*\* amyloid-peptide-induced, basal forebrain (BF) lesioned and carbon monoxide (CO)-induced amnesia models and senescence-accelerated mouse (SAM). In addition, these \*\*\*sigma1\*\*\* - \*\*\*receptor\*\*\* agonists have good profiles for the central acetylcholine and dopamine systems. Moreover, they also have neuroprotective and anti-depressive effects. These evidence suggested that the \*\*\*sigma1\*\*\* - \*\*\*receptor\*\*\* agonists may be promising compounds for the treatment of dementing disorders such as Alzheimer's disease, senile dementia and vascular dementia. However, the sigma-receptor family is still considered to be enigmatic molecular targets. More molecular cloning and biochemical studies on the sigma-receptor family are needed.

L3 ANSWER 21 OF 21 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.  
on STN  
ACCESSION NUMBER: 94382870 EMBASE  
DOCUMENT NUMBER: 1994382870  
TITLE: Selective antagonism of opioid analgesia by a sigma system.  
AUTHOR: Chien C.-C.; Pasternak G.W.  
CORPORATE SOURCE: Department of Neurology, Memorial Sloan-Kettering Cancer Ctr., 1275 York Avenue, New York, NY 10021, United States  
SOURCE: Journal of Pharmacology and Experimental Therapeutics, (1994) 271/3 (1583-1590).  
ISSN: 0022-3565 CODEN: JPETAB  
COUNTRY: United States  
DOCUMENT TYPE: Journal; Article  
FILE SEGMENT: 008 Neurology and Neurosurgery  
030 Pharmacology  
037 Drug Literature Index  
LANGUAGE: English  
SUMMARY LANGUAGE: English  
AB (+)Pentazocine antagonizes morphine analgesia as potently as its (-) isomer, ruling out an opioid receptor mechanism of action and suggesting, which suggests a role for \*\*\*sigma1\*\*\* \*\*\*receptors\*\*\* . Systemic (+) pentazocine also reverses supraspinal or spinal morphine analgesia. 1,3-Di(2-tolyl)guanidine, a sigma ligand with no appreciable opioid receptor affinity, antagonizes morphine analgesia. The actions of both (+)pentazocine and 1,3-di(2- tolyl)guanidine are reversed by haloperidol, which has high affinity for both sigma and D2 receptors, but not by the D2-selective antagonist (-)sulpiride, which lacks activity at sigma sites. The antiopioid sigma system is tonically active. Haloperidol, but not (-)sulpiride, decreases morphine ED50 almost 2-fold. The antiopioid system modulates only mu analgesia. Unlike analgesia, (+)pentazocine does not influence morphine's inhibition of gastrointestinal transit or

lethality. (+)Pentazocine also antagonizes kappa1, kappa3 and delta analgesia through sigma mechanisms in a haloperidol-sensitive manner. (-)Sulpiride is inactive. Alone, haloperidol enhances kappa1, kappa3 and delta analgesia more dramatically than morphine, which indicates that the sigma system is active against all opioid analgesic systems. Sigma systems are responsible for some strain differences in kappa receptor sensitivity. Unlike CD-1 mice, BALB-C mice are relatively insensitive toward the kappa1 agent U50,488H and the kappa3 analgesic naloxone benzoylhydrazone. Blockade of the sigma system with haloperidol eliminates these strain differences. In conclusion, sigma1 systems functionally antagonize opioid analgesia without affecting morphine's effects on gastrointestinal transit or lethality. The antiopioid sigma system is tonically active and is more active against kappa analgesia than mu. The level of this tonic activity plays a significant role in strain differences in analgesic sensitivity toward opioid analgesia.